Differential Effects of Midazolam and Zolpidem on Sleep-Wake States and Epileptic Activity in WAG/Rij Rats

H. DEPOORTERE,* D. FRANCON,* E. L. J. M. VAN LUIJTELAAR,† W. H. I. M. DRINKENBURG† AND A. M. L. COENEN†

*Synthélabo Recherche (L.E.R.S.), 31, avenue Paul Vaillant Couturier, F-92220 Bagneux, France
†NICI, Department of Psychology, University of Nijmegen, Nijmegen, The Netherlands

Received 1 September 1994; Revised 14 January 1995; Accepted 17 January 1995

DEPOORTERE, H., D. FRANCON, E. L. J. M. VAN LUIJTELAAR, W. H. I. M. DRINKENBURG AND A. M. L. COENEN. Differential effects of midazolam and zolpidem on sleep-wake states and epileptic activity in WAG/Rij rats. PHARMACOL BIOCHEM BEHAV 51(4) 571-576, 1995. — Hypnotic drugs are known to possess antiepileptic activity. Therefore, the effects of the benzodiazepine hypnotic midazolam (10 mg/kg) and the novel imidazopyridine hypnotic zolpidem (10 mg/kg) on sleep-wake states and on the number of spike-wave discharges were evaluated in WAG/Rij rats. Rats of this strain are considered to be a model for generalized absence epilepsy. Animals were implanted with chronic monopolar EEG electrodes and, after recovery from surgery, the EEG was recorded for 6 h during the dark period on 3 consecutive days. Sleep recordings were analyzed using Hjorth's parameters and number and duration of spike-wave discharges were visually determined. It was found that both drugs facilitated nonREM sleep at the cost of wakefulness. Both hypnotics also reduced the number and duration of spike-wave discharges. The initial decrease after midazolam, however, was followed by a rebound reflecting a poorer quality of vigilance expressed as an increase in spike-wave discharges. The strong antiabsence activity of zolpidem mimics that of midazolam and is well correlated with their equipotent hypnotic action and anticonvulsant effect in the isoniazid test.

Sleep-wake states  Spike-wave discharges  Absence epilepsy  Midazolam  Zolpidem  Hypnotics
WAG/Rij strain  Rats  Rebound effects

FOR THE last 2 decades benzodiazepines have been the anxio-
ytics and hypnotics of first choice due to their efficacy, their
relatively mild side effects, and their safety. A standing prob-
lem is that they also induce daytime drowsiness, which is often
accompanied by muscular relaxation (6). Moreover, it has be-
come increasingly clear that benzodiazepines affect cognitive
functioning and psychomotor performance (16,20,25). Next
to all these actions, benzodiazepines show broad spectrum
antiepileptic properties: they reliably reduce various types of
convulsions and counteract spike-wave discharges.

Until recently, it was generally assumed that these different
effects of benzodiazepines are indissolubly linked. Nowadays,
research efforts are aimed at separating the different actions
of the benzodiazepines, and new drugs are currently being
developed that do not possess all the effects of the benzodia-
zepines. Investigations of partial agonists at the benzodiazepine
(ω) receptors suggest that it is, indeed, possible to separate
these effects. Concrete examples were presented by Coenen
and van Luijteelaar (3), who demonstrated that the hypnotic
and antiepileptic action of ZK 91296 were discriminable.
Stephens et al. (24) and Coenen et al. (5) also showed that the
anxiolytic and hypnotic actions of abecarnil were distinguish-
able.

The novel hypnotic, zolpidem, an imidazopyridine deriva-
tive, shares many of the pharmacological actions characteris-
tic of benzodiazepines. There are, however, differences such
as the weaker myorelaxant effects of zolpidem (1,10). De-
poortere et al. (10) investigated the anticonvulsant profile
of zolpidem in pentylenetetrazol-induced convulsions and in
the electroshock model. They demonstrated a considerable
weaker anticonvulsant activity for zolpidem than for the ben-
zodiazepines midazolam, triazolam, and flunitrazepam in

1 To whom requests for reprints should be addressed.
these models. However both categories of drugs, ligands of the GABA-benzodiazepine receptor complex, have equipotent hypnotic activity (11), as well as anticonvulsant effects in the isoniazid convulsant model (30). Based on the latter findings, it is likely that zolpidem, like the benzodiazepines, inhibits spike-wave discharges.

The purpose of the present study was to evaluate the effects on sleep-wake states and on the amount of epileptic activity of the classical benzodiazepine hypnotic midazolam and the nonbenzodiazepine hypnotic zolpidem. This was done in WAG/Rij rats, which show spontaneously occurring spike-wave discharges in the EEG and are considered as an adequate model for human absence epilepsy (4,27).

METHOD

Experiments were performed on 10 male WAG/Rij rats, weighing between 320 and 400 g. They were maintained under identical environmental conditions, under a 12 L:12 D cycle and a constant temperature of 21°C. Rats were anesthetized with an intraperitoneal injection of 50 mg/kg sodium pentobarbital and implanted with three small electrode screws on each hemisphere. The first electrode was screwed into the skull above the sensorimotor cortex (1.5 mm lateral to the median suture and 1.5 mm behind the fronto-parietal suture), the second one into the skull above the visual cortex (1.5 mm lateral to the median suture and 1.5 mm in front of the parieto-occipital suture), and the third one above the cerebellum. The latter served as a reference electrode. All electrodes were attached to a seven-lead Winchester connector fixed with dental cement to the cranium. Three weeks of postoperative recovery were allowed. During this period, half of the rats were habituated to a reversed light: dark cycle, whereas the other half kept the regular cycle with lights on between 0700 and 1900 h. There was no difference between the two groups of five rats: they had the same ages and weights (360 ± 20 g). Four days before the experiment, the animals were placed in Plexiglas cylinders with free access to food and water (habituation period to experimental conditions). Each rat was injected twice at intervals of at least 3 weeks and recorded from during the light period after treatment with vehicle (control states in normal rats (9-11). The other five rats were recorded after treatment with drug (either zolpidem or midazolam because of their equal potencies on sleep-wake states in normal rats (9-11)). One day later, non-REM sleep was reduced and wakefulness was increased for the first hour of recording. Zolpidem decreased wakefulness and increased non-REM sleep during the first 2 h following administration. No significant effects were seen on REM sleep. The data are shown in Fig. 1a. One day later, non-REM sleep was reduced and wakefulness was increased for the first hour of recording. Zolpidem decreased the number and total duration of spike-wave discharges (Fig. 2a). These effects were present during the first 3 h after administration. The number and duration of the spike-wave discharges on the postdrug day did not differ from predrug values, except that the duration was still reduced during the first hour on the postdrug day.

Midazolam increased non-REM sleep and reduced wakefulness. The results are shown in Fig. 1b. Midazolam reduced the number and duration of spike-wave discharges during the first 3 h following drug administration (Fig. 2b). This initial decrease was followed by a significant increase in number and total duration at the fifth hour of the experimental day. The total duration of the spike-wave discharges was still enhanced.

The EEG was automatically analyzed by a computerized system that discriminates between these states. The system, which uses Hjorth's descriptors, has been described in detail, validated, and extensively used (8,15). The performance of the system was regularly checked against visual scoring. Baseline parameters for the light and dark period were determined, while drug effects were only evaluated in the dark period, the most sensitive part of the light: dark cycle to evaluate hypnotic effects and antilapse effect on spike-wave discharges (9,28,29). Spike-wave discharges were counted based on criteria published elsewhere, whereby the number and total duration of the spike-wave discharges per hour were determined (27). Within-group comparisons were made with Student's t-test for paired observations.

RESULTS

Sleep-wakefulness characteristics for the baseline light and dark period are presented in Table 1. This table shows that the rats predominantly slept during the light period (74% of total time) and that the percentages of non-REM sleep and REM sleep, the number of REM periods, and the mean length of periods of REM sleep were greater in the light than in the dark period. The rats had more spike-wave discharges during the dark period (mean and standard error: 197 ± 21) than during the light period (85 ± 8).

Zolpidem reduced wakefulness and increased non-REM sleep during the first 2 h following administration. No significant effects were seen on REM sleep. The data are shown in Fig. 1a. One day later, non-REM sleep was reduced and wakefulness was increased for the first hour of recording.

Zolpidem decreased the number and total duration of spike-wave discharges (Fig. 2a). These effects were present during the first 3 h after administration. The number and duration of the spike-wave discharges on the postdrug day did not differ from predrug values, except that the duration was still reduced during the first hour on the postdrug day.

Midazolam increased non-REM sleep and reduced wakefulness. The results are shown in Fig. 1b. Midazolam reduced the number and duration of spike-wave discharges during the first 3 h following drug administration (Fig. 2b). This initial decrease was followed by a significant increase in number and total duration at the fifth hour of the experimental day. The total duration of the spike-wave discharges was still enhanced.

TABLE 1

<table>
<thead>
<tr>
<th>Recordings During</th>
<th>W% ± SEM</th>
<th>non-REM% ± SEM</th>
<th>REM% ± SEM</th>
<th>nREM ± SEM</th>
<th>tREM s ± SEM</th>
<th>nSWD ± SEM</th>
<th>dSWD s ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Light</td>
<td>24.2 ± 1.5</td>
<td>65.4 ± 1.1</td>
<td>10.4 ± 1.3</td>
<td>19 ± 1</td>
<td>116 ± 7</td>
<td>85 ± 8</td>
<td>440 ± 82</td>
</tr>
<tr>
<td>Dark</td>
<td>59.2 ± 2.5</td>
<td>38.6 ± 2.5</td>
<td>2.2 ± 0.8</td>
<td>7 ± 2</td>
<td>89 ± 3</td>
<td>197 ± 21</td>
<td>1534 ± 159</td>
</tr>
</tbody>
</table>

The results (± SEM) are expressed as percentages for wakefulness (W), non-REM sleep, REM sleep, number of REM periods (nREM), mean duration of REM periods in s (tREM), number of spike-wave discharges (nSWD), and total duration of spike-wave discharges in seconds (dSWD) in a 6-h recording period during the first control day.
FIG. 1. (a) Effects of zolpidem (10 mg/kg PO) on sleep-wake states in WAG/Rij rats recorded during the dark period (n = 5). *p ≤ 0.05; **p ≤ 0.01 vs. control values. (b) Effects of midazolam (10 mg/kg PO) on sleep-wake states in WAG/Rij rats recorded during the dark period (n = 5). *p ≤ 0.05; **p ≤ 0.01 vs. control values.

An overview of the main outcomes is presented in Table 2. Both drugs facilitated non-REM sleep at the expense of wakefulness and reduced the number and duration of spike-wave discharges with a similar efficacy. On the second control day, the two drugs differed in that zolpidem decreased non-REM sleep and the number of spike-wave discharges while midazolam increased spike-wave activity.

DISCUSSION

The sleep-wakefulness profile of WAG/Rij rats differs from that commonly seen in other laboratory rats. The low percentage REM sleep for the dark period (2%) seems to be smaller than commonly reported (5%) (2,9,18,26). Also, the duration of the REM periods seems to be shorter than found earlier for outbreds in the same laboratory (7). Thus, it seems that WAG/Rij rats spend less time in REM sleep than other rats, and this was directly evaluated by Gandolfo et al. (13) who reported that WAG/Rij rats have less REM sleep periods per hour, with a lower percentage of REM sleep, and that these rats seemed to have difficulties entering into REM sleep.

Despite differences in genotype between outbred and WAG/Rij rats, it seems that both midazolam and zolpidem reliably induce nonREM sleep at the cost of wakefulness (Table 2). The hypnotic effects of zolpidem were earlier described by Depoortere et al. (10,11), and the present results suggest that the hypnotic effects of both drugs are independent of the rat's genotype. However, the two drugs differ in their effects on rebound spike-wave discharges and on sleep 24 h following...
FIG. 2. (a) Effects of zolpidem (10 mg/kg PO) on number and total duration of spike-wave discharges in WAG/Rij rats (n = 5). *p ≤ 0.05; **p ≤ 0.01 vs. control values. (b) Effects of midazolam (10 mg/kg PO) on number and total duration of spike-wave discharges in WAG/Rij rats (n = 5). *p ≤ 0.05; **p ≤ 0.01 vs. control values.
EFFECTS OF MIDAZOLAM AND ZOLPIDEM IN WAG/Rij RATS

The decrease in the number and duration of spike-wave discharges after midazolam was expected considering the well-known antiepileptic action of benzodiazepines (22). However, the reduction in spike-wave discharges seen with zolpidem was as marked as that of midazolam. It is known that the anticonvulsant action of zolpidem in pentylentetrazol-induced seizures or in electroshock convulsions is less than those of the benzodiazepines flunitrazepam and midazolam (10). Thus, zolpidem is 225 and 69 times less potent than flunitrazepam and midazolam, respectively, in pentylenetetrazol-induced convulsions, and 13 and 8 times less potent in electroshock convulsions. On the other hand, zolpidem is more active in convulsive models that involve impairment in GABAergic transmission, e.g., isoniazid-induced convulsions. It possesses the same or an even higher efficacy as compared to flunitrazepam and midazolam, in this model (30). The present data shows that midazolam and zolpidem are equally effective in suppressing spike-wave discharges after midazolam was followed by an increase in the number of spike-wave discharges. To our knowledge, rebound effects on spontaneous epileptic activity have not previously been reported. Only following abrupt cessation of chronic benzodiazepine administration was an increase in susceptibility for epileptic seizures reported (23). The rebound effects seen only on spike-wave discharges but not on sleep after midazolam could reflect a poorer quality of vigilance, with an increase of quiet wakefulness–drowsiness periods, because spike-wave discharges preferentially occur during periods with a low level of vigilance (12). In contrast, zolpidem maintains an adequate level of vigilance after its hypnotic effect during the waking (dark) period without rebound of spike-wave discharges.

In all, the present data show comparable hypnotic effects of zolpidem and midazolam in rats with a different genotype than commonly used. They further suggest an equal potency of these drugs to suppress spike-wave discharges in WAG/Rij rats. Finally, the data show a rebound effect on spike-wave activity for midazolam that could reflect a poorer quality of vigilance, but this warrants further investigation.

ACKNOWLEDGEMENTS

The authors gratefully acknowledge A. Clemente and C. Laufrais for their skilled experimental assistance and 1. Chartouni for her secretarial help.

REFERENCES